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FILE COVERS 1907 - 29 Jul 2009 VOL 151 ISS 5
FILE LAST UPDATED: 28 Jul 2009 (20090728/ED)
REVISED CLASS FIELDS (/NCL) LAST RELOADED: Jun 2009
USPTO MANUAL OF CLASSIFICATIONS THESAURUS ISSUE DATE: Jun 2009

CAplus now includes complete International Patent Classification (IPC) reclassification data for the second quarter of 2009.

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=>

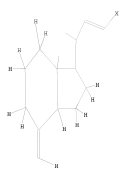
Uploading C:\Program Files\Stnexp\Queries\10579594.str

L1 STRUCTURE UPLOADED

=> d

L1 HAS NO ANSWERS

L1 STR



Structure attributes must be viewed using STN Express query preparation.

=> s 11 sss sam

REG1stRY INITIATED

Substance data SEARCH and crossover from CAS REGISTRY in progress... Use DISPLAY HITSTR (or FHITSTR) to directly view retrieved structures.

SAMPLE SEARCH INITIATED 13:48:30 FILE 'REGISTRY'
SAMPLE SCREEN SEARCH COMPLETED - 16 TO ITERATI

100.0% PROCESSED 16 ITERATIONS SEARCH TIME: 00.00.01

1 ANSWERS

FULL FILE PROJECTIONS: ONLINE **COMPLETE**
BATCH **COMPLETE**
PROJECTED ITERATIONS: 80 TO 560
PROJECTED ANSWERS: 1 TO 80

L2 1 SEA SSS SAM L1

L3 1 L2

=> d ibib abs hitstr THE ESTIMATED COST FOR THIS REQUEST IS 5.64 U.S. DOLLARS

DO YOU WANT TO CONTINUE WITH THIS REQUEST? (Y) /N:v

ANSWER 1 OF 1 CAPLUS COPYRIGHT 2009 ACS on STN

ACCESSION NUMBER: 2004:1024083 CAPLUS

DOCUMENT NUMBER: 142:134781

TITLE: Potent, Selective and Low-Calcemic Inhibitors of CYP24 Hydroxylase: 24-Sulfoximine Analogues of the Hormone

10,25-Dihydroxyvitamin D3

AUTHOR(S): Kahraman, Mehmet; Sinishtaj, Sandra; Dolan, Patrick M.; Kensler, Thomas W.; Peleg, Sara; Saha, Uttam;

Chuang, Samuel S.; Bernstein, Galina; Korczak, Bozena; Posner, Garv H.

CORPORATE SOURCE: Department of Chemistry, School of Arts and Sciences,

The Johns Hopkins University, Baltimore, MD, 21218,

SOURCE: Journal of Medicinal Chemistry (2004), 47(27),

6854-6863

CODEN: JMCMAR; ISSN: 0022-2623 PUBLISHER: American Chemical Society

т

DOCUMENT TYPE: Journal

LANGUAGE: English OTHER SOURCE(S): CASREACT 142:134781

AB A dozen 24-sulfoximine analogs of the hormone 1a, 25-dihydroxyvitamin D3 were prepared, differing not only at the stereogenic sulfoximine stereocenter but also at the A-ring. Although these sulfoximines were not active transcriptionally and were only very weakly antiproliferative, some of them are powerful hydroxylase enzyme inhibitors. Specifically, 24(S)-NH Ph sulfoximine I is an extremely potent CYP24 inhibitor (IC50 = 7.4 nM) having low calcemic activity. In addition, this compound shows high selectivity toward the CYP24 enzyme in comparison to CYP27A1 (IC50 > 1000 nM) and CYP27B (IC50 = 554 nM).

825638-30-0P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation and CYP24 inhibitory activity of dihydroxyvitamin D3 sulfoximine analogs)

RN 825638-30-0 CAPLUS

CN Silanamine, N-[(R)-[(1E,3S)-3-[(1R,3aS,4E,7aR)-4-[(2Z)-2-[(3S,5R)-3,5-bis[[(1,1-dimethylethyl)dimethylsilyl]oxy]-2-

methylenecyclohexylidene]ethylidene]octahydro-7a-methyl-1H-inden-1-yl]-1-fluoro-1-buten-1-yl]oxidophenyl-\lambda-sulfanylidene]-1-(1,1-dimethyl-1,0-

Absolute stereochemistry.

Double bond geometry as shown.

OS.CITING REF COUNT: 27 THERE ARE 27 CAPLUS RECORDS THAT CITE THIS

RECORD (27 CITINGS)
REFERENCE COUNT: 36 THERE ARE 36 CITED REFERENCES AVAILABLE FO

THERE ARE 36 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

=> s l1 sss full

REG1stRY INITIATED

Substance data SEARCH and crossover from CAS REGISTRY in progress...
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FULL SEARCH INITIATED 13:49:31 FILE 'REGISTRY'

FULL SCREEN SEARCH COMPLETED - 268 TO ITERATE

100.0% PROCESSED 268 ITERATIONS SEARCH TIME: 00.00.01 16 ANSWERS

L4 16 SEA SSS FUL L1

L5 7 L4

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L6 2 L5 AND PY<2003

=> s 15 and py<2004 24035992 PY<2004

L7 2 L5 AND PY<2004

=> d 1-2 ibib abs hitstr

THE ESTIMATED COST FOR THIS REQUEST IS 11.28 U.S. DOLLARS DO YOU WANT TO CONTINUE WITH THIS REQUEST? (Y)/N:y

L7 ANSWER 1 OF 2 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 1994:245588 CAPLUS

DOCUMENT NUMBER: 120:245588

ORIGINAL REFERENCE NO.: 120:43561a,43564a

TITLE: 1\(\alpha\), 24S-Dihydroxy-26,27-cyclo-22-yne vitamin D3: the side chain triple bond analog of MC 903

(calcipotriol)

AUTHOR(S): Calverley, Martin J.: Bretting, Claus Aa.S.

AUTHOR(S): Calverley, Martin J.; Bretting, Claus Aa.S.
CORPORATE SOURCE: Chem. Res. Dep., Leo Pharm. Prod., Ballerup, DK-2750,

Den.

SOURCE: Bioorganic & Medicinal Chemistry Letters (1993

), 3(9), 1841-4

CODEN: BMCLE8; ISSN: 0960-894X PE: Journal

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 120:245588

GI

The side chain propargylic alc. function [established stereoselectively AR via S-Alpine-Borane reduction of ynone I (TBDMS = tert-butyldimethylsilyl) and correlated with MC 903] in the title compound II replaces the metabolically labile allylic alc. function of MC 903, a selective analog of the vitamin D hormone used for treating psoriasis. II exhibits reduced in vitro activity but still shows selectively much lower in vivo calcemic effects.

154171-12-7P RL: SPN (Synthetic preparation); PREP (Preparation)

(preparation and lithiation and cyclopropylcarbonylation of)

RN 154171-12-7 CAPLUS CN

Silane, [[(1α,3β,5E,7E)-23,23-dichloro-24-nor-9,10-secochola-5,7,10(19),22-tetraene-1,3-diyl]bis(oxy)]bis[(1,1-dimethylethyl)dimethyl-(9CI) (CA INDEX NAME)

Absolute stereochemistry. Double bond geometry as shown.

L7 ANSWER 2 OF 2 CAPLUS COPYRIGHT 2009 ACS on STN

ACCESSION NUMBER: 1992:255875 CAPLUS DOCUMENT NUMBER: 116:255875

ORIGINAL REFERENCE NO.: 116:43403a,43406a

TITLE:

Preparation of vitamin D analogs as drugs Bretting, Claus Aage Svensgaard INVENTOR(S):

PATENT ASSIGNEE(S): Leo Pharmaceutical Products Ltd. A/S. Den.

SOURCE: PCT Int. Appl., 40 pp.

CODEN: PIXXD2 DOCUMENT TYPE: Patent

English LANGUAGE: FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.					KIND DATE		APPLICATION NO.				DATE						
WO	WO 9203414				A1		19920305			WO 1991-DK200					19910711 <		
	W:	ΑU,	BB,	BG,	BR,	CA,	CS,	FΙ,	HU,	JP,	KP,	KR,	LK,	MC,	MG,	MN,	MW,
					SD,												
	RW:									CM,		DK,	ES,	FR,	GA,	GB,	GN,
		GR,	IT,	LU,	ML,	MR,	NL,	SE,	SN,	TD,	TG						
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	EP	543864			B1	19941214					
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	FI	103791			B1	19990930					
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	LV	10089			В	19941020	LV	1993-243		19930215	<
	LT	3666			В	19960125	LT	1993-965		19930910	<
PRIC	RITY	APPLN	. IN	FO.:			GB	1990-17890	A	19900815	
							CS	1992-3726	A	19910711	
							WO	1991-DK200	A	19910711	

OTHER SOURCE(S):

MARPAT 116:255875

AB Title compds. [I; R = ZIC.tplbond.CZZCRIRZX; Rl, R2 = H, hydrocarbyl; or R1R2 = atoms to form a carbocyclic ring; R3 = cyclohexylidenemethylidyne group Q1; X = H, OH; Z1 = (substituted)(CR2)m; Z2 = bond, hydrocarbylenediyl; m = 0-2] were prepared as antiinflammatories, immunomodulators, etc. (no data). Thus, I (R = CHO, R3 = cyclohexylidenemethylidyne group Q2) was condensed with (Me2N)3P:CCL2 (prepared in situ) and the product treated, in turn, with BuLi and Br(CR2)3CBt2OS1Me3 to give I [R = C.tplbond.C(CR2)3CBt2OS1Me3, R3 = Q2] which was photoisomerized to give, after deprotection, I [R = C.tplbond.C(CR2)3CBt2OR, R3 = Q1]

IT 141545-84-8P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation and reaction of, in preparation of antiinflammatory and immunomodulator)

RN 141545-84-8 CAPLUS

CN 1H-Indene, 4-[(2E)-2-[(3S,5R)-3,5-bis[[(1,1-

4

 $\label{limit} $$ \dim \theta_1=1-[(1R)-3,3-dichloro-1-methyl-2-propen-1-yl] octahydro-7a-methyl-, $$$

(1R, 3aS, 4E, 7aR) - (CA INDEX NAME)

OS.CITING REF COUNT:

9 THERE ARE 9 CAPLUS RECORDS THAT CITE THIS RECORD (9 CITINGS)

REFERENCE COUNT:

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